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[General] [Name and origin] [References] [Comments] [Cross-references] [Keywords] [Features] [Sequence] [Tools]

General information about the entry

Entry name

SP3E BACSU

Primary accession number

P21458

Secondary accession number

P21459

Entered in SWISS-PROT in

Release 18, May 1991

Sequence was last modified in

Release 31, February 1995

Annotations were last modified in

Release 41, June 2002

Name and origin of the protein

Protein name

Stage III sporulation protein E

Synonyms

None

Gene name

SPOIIIE

From

Bacillus subtilis [TaxID: 1423]

Taxonomy

Bacteria; Firmicutes; Bacillales; Bacillaceae; Bacillus.

References

[1] SEQUENCE FROM NUCLEIC ACID.

STRAIN=168:

MEDLINE=88199510; PubMed=3129532; [NCBI, ExPASy, EBI, Israel, Japan]

Butler P.D., Mandelstam J.;

"Nucleotide sequence of the sporulation operon, spoIIIE, of Bacillus subtilis.";

J. Gen. Microbiol. 133:2359-2370(1987).

[2] REVISIONS.

MEDLINE=90014185; PubMed=2507870; [NCBI, ExPASy, EBI, Israel, Japan]

Foulger D., Errington J.;

"The role of the sporulation gene spoIIIE in the regulation of prespore-specific gene expression in Bacillus subtilis.";

Mol. Microbiol. 3:1247-1255(1989).

[3]

SEQUENCE FROM NUCLEIC ACID.

STRAIN=168:

MEDLINE=98044033; PubMed=9384377; [NCBI, ExPASy, EBI, Israel, Japan]

Kunst F., Ogasawara N., Moszer I., Albertini A.M., Alloni G., Azevedo V., Bertero M.G., Bessieres P., Bolotin A., Borchert S., Borriss R., Boursier L., Brans A., Braun M., Brignell S.C., Bron S.,

Brouillet S., Bruschi C.V., Caldwell B., Capuano V., Carter N.M., Choi S.K., Codani J.J.,

Connerton I.F., Cummings N.J., Daniel R.A., Denizot F., Devine K.M., Dusterhoft A., Ehrlich S.D.,

Emmerson P.T., Entian K.D., Errington J., Fabret C., Ferrari E., Foulger D., Fritz C., Fujita M., Fujita Y., Fuma S., Galizzi A., Galleron N., Ghim S.Y., Glaser P., Goffeau A., Golightly E.J., Grandi G., Guiseppi G., Guy B.J., Haga K., Haiech J., Harwood C.R., Henaut A., Hilbert H., Holsappel S., Hosono S., Hullo M.F., Itaya M., Jones L., Joris B., Karamata D., Kasahara Y., Klaerr-Blanchard M., Klein C., Kobayashi Y., Koetter P., Koningstein G., Krogh S., Kumano M., Kurita K., Lapidus A., Lardinois S., Lauber J., Lazarevic V., Lee S.M., Levine A., Liu H., Masuda S., Mauel C., Medigue C., Medina N., Mellado R.P., Mizuno M., Moestl D., Nakai S., Noback M., Noone D., O'Reilly M., Ogawa K., Ogiwara A., Oudega B., Park S.H., Parro V., Pohl T.M., Portetelle D., Porwollik S., Prescott A.M., Presecan E., Pujic P., Purnelle B., Rapoport G., Rey M., Reynolds S., Rieger M., Rivolta C., Rocha E., Roche B., Rose M., Sadaie Y., Sato T., Scanlan E., Schleich S., Schroeter R., Scoffone F., Sekiguchi J., Sekowska A., Seror S.J., Serror P., Shin B.S., Soldo B., Sorokin A., Tacconi E., Takagi T., Takahashi H., Takemaru K., Takeuchi M., Tamakoshi A., Tanaka T., Terpstra P., Tognoni A., Tosato V., Uchiyama S., Vandenbol M., Vannier F., Vassarotti A., Viari A., Wambutt R., Wedler E., Wedler H., Weitzenegger T., Winters P., Wipat A., Yamamoto H., Yamane K., Yasumoto K., Yata K., Yoshida K., Yoshikawa H.F., Zumstein E., Yoshikawa H., Danchin A.;

"The complete genome sequence of the Gram-positive bacterium Bacillus subtilis."; Nature 390:249-256(1997).

[4] CHARACTERIZATION.

MEDLINE=94212172; PubMed=8160014; [NCBI, ExPASy, EBI, Israel, Japan] Wu L.J., Errington J.;

"Bacillus subtilis spoIIIE protein required for DNA segregation during asymmetric cell division."; Science 264:572-575(1994).

Comments

- FUNCTION: REQUIRED FOR SPORULATION. IT IS REQUIRED FOR CHROMOSOME SEGREGATION INTO THE PRESPORE COMPARTMENT. MAY ALSO BE REQUIRED TO PREVENT SIGMA F FROM BECOMING ACTIVE IN THE MOTHER CELL. MAY BIND DNA.
- SUBCELLULAR LOCATION: Integral membrane protein (Potential).
- SIMILARITY: BELONGS TO THE FTSK/SPOILIE FAMILY.

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Cross-references

M17445; AAA22784.1; ALT_SEQ.[EMBL / GenBank / DDBJ] [CoDingSequence]

EMBL M17445; AAA22785.1; ALT_SEQ.[EMBL / GenBank / DDBJ] [CoDingSequence]

Z99112; CAB13553.1; -. [EMBL / GenBank / DDBJ] [CoDingSequence]

A32269; A32269.

PIR B32269; B32269.

S09411; S09411.

SubtiList BG10763; spoIIIE. [SubtiList / NRSub / Micado]

InterPro IPR002543; FtsK_SpoIIIE.

Graphical view of domain structure.

Pfam <u>PF01580;</u> FtsK SpoIIIE; 1.

ProDom [Domain structure / List of seq. sharing at least 1 domain].

BLOCKS <u>P21458</u>. ProtoNet <u>P21458</u>.

 ProtoMap
 P21458.

 PRESAGE
 P21458.

 DIP
 P21458.

 ModBase
 P21458.

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Keywords

Sporulation; ATP-binding; Transmembrane; Complete proteome.

Features

Key	From	To I	Length	Description
TRANSMEM	22	42	21	POTENTIAL.
TRANSMEM	52	72	21	POTENTIAL.
TRANSMEM	87	107	21	POTENTIAL.
TRANSMEM	154	1/4	21	POTENTIAL.
NP BIND	467	474	8	ATP (POTENTIAL).



Feature aligner



Feature table viewer

Sequence information

Length: 787 AA	Molecular v Da	weight: 87153	CRC64: 310 sequence]	6F40F611 28 1	6E3 [This is a	checksum on the
10	20	30	40	50	60	
 MAKKKRKSRK	KOAKOTNIKA 	ELNGLLCIAI	 SIIAILQLGV	 VGQTFIYLFR	 FFAGEWFILC	
7 Ó	80	90	100	110	120	
l LLGLLVLGVS	LFWKKKTPSL	LTRRKAGLYC	IIASILLLSH	VQLFKNLTHK	GSİESASVVR	
130	140	150	160	170	180	
 NTWELFLMDM	NGSSASPDLG	GGMIGALLFA	ASHFLFASTG	SQIMAIVMIL	 IGMILVTGRS	
190	200	210	220	230	240	
LQETLKKWMS	PIGRFIKEQW	LAFIDDMKSF	KSNMQSSKKT	 KAPSKKQKPA	RKKQQMEPEP	
250	260	270	280	290	300	
 PDEEGDYETV	SPLIHSEPII	 SSFSDRNEEE	 ESPVIEKRAE	PVSKPLQDIQ 	 PETGDQETVS	
310	320	330	340	350	360	
 APPMTFTELE	 NKDYEMPSLD	 LLADPKHTGQ	QADKKNIYEN	 ARKLERTFQS	 FGVKAKVTQV	
370 	380	390 I	400	410	420	
HLGPAVTKYE	VYPDVGVKVS	KIVNLSDDLA	LALAAKDIRI	EAPIPGKSAI	GIEVPNAEVA	
430	440	450	460	470	480	
MVSLKEVLES	KLNDRPDANV	LIGLGRNISG	EAVLAELNKM	PHLLVAGATG	SGKSVCVNGI	
490 	500 I	510 	520 I	530 	540 I	

ITSILMRAKP	HEVKMMMIDP	KMVELNVYNG	IPHLLAPVVT	DPKKASQALK	KVVNEMERRY
550	560	570	580	590	600
ELFSHTGTRN	IEGYNDYIKR	ANNEEGAKQP	ELPYIVVIVD	ELADLMMVAS	SDVEDSITRL
610	620	630	640	650	660
SQMARAAGIH	LIIATQRPSV	DVITGVIKAN	IPSRIAFSVS	SQTDSRTILD	MGGAEKLLGR
670	680	690	700	710	720
			I		I
GDMLFLPVGA	NKPVRVQGAF	LSDDEVEKVV	DHVITQQKAQ	YQEEMIPEET	TETHSEVTDE
730	740	750	760	770	780
	I				
LYDEAVELIV	GMQTASVSML	QRRFRIGYTR	AARLIDAMEE	RGVVGPYEGS	KPREVLLSKE

KYDELSS

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Direct BLAST submission at NCBI (Bethesda, <u>USA)</u>



ScanProsite, MotifScan



Sequence analysis tools: <u>ProtParam</u>, <u>ProtScale</u>, Compute pI/Mw, PeptideMass, PeptideCutter, Dotlet (Java)



Feature table viewer (Java)



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[General] [Name and origin] [References] [Comments] [Cross-references] [Keywords] [Features] [Sequence] [Tools]

General information about the entry

Entry name

FTSI ECOLI

Primary accession number

P04286

Secondary accession numbers

None

Entered in SWISS-PROT in

Release 04, March 1987

Sequence was last modified in

Release 04, March 1987 Release 41, June 2002

Annotations were last modified in

Name and origin of the protein

Peptidoglycan synthetase ftsI [Precursor]

Protein name Synonyms

Penicillin-binding protein 3

PBP-3

Gene name

FTSI or PBPB or <u>B0084</u> or <u>Z0094</u> or <u>ECS0088</u>

Bacteria; Proteobacteria; Gammaproteobacteria;

From

Escherichia coli

[TaxID: <u>5</u>62]

Escherichia coli O157:H7 [TaxID: 83334]

Taxonomy

Enterobacteriaceae; Escherichia.

References

[1] SEQUENCE FROM NUCLEIC ACID.

STRAIN=K12;

MEDLINE=83296957; PubMed=6350821; [NCBI, ExPASy, EBI, Israel, Japan]

Nakamura M., Maruyama I.N., Soma M., Kato J., Suzuki H., Horota Y.;

"On the process of cellular division in Escherichia coli: nucleotide sequence of the gene for penicillin-binding protein 3.";

Mol. Gen. Genet. 191:1-9(1983).

[2] SEQUENCE FROM NUCLEIC ACID.

STRAIN=K12;

MEDLINE=92334977; PubMed=1630901; [NCBI, ExPASy, EBI, Israel, Japan]

Yura T., Mori H., Nagai H., Nagata T., Ishihama A., Fujita N., Isono K., Mizobuchi K., Nakata A.; "Systematic sequencing of the Escherichia coli genome: analysis of the 0-2.4 min region."; Nucleic Acids Res. 20:3305-3308(1992).

[3]

SEQUENCE FROM NUCLEIC ACID.

STRAIN=K12 / MG1655;

MEDLINE=97426617; PubMed=9278503; [NCBI, ExPASy, EBI, Israel, Japan]

Blattner F.R., Plunkett G. III, Bloch C.A., Perna N.T., Burland V., Riley M., Collado-Vides J., Glasner J.D., Rode C.K., Mayhew G.F., Gregor J., Davis N.W., Kirkpatrick H.A., Goeden M.A., Rose D.J., Mau B., Shao Y.;

"The complete genome sequence of Escherichia coli K-12."; Science 277:1453-1474(1997).

[4] SEQUENCE FROM NUCLEIC ACID.

STRAIN=0157:H7 / EDL933 / ATCC 700927;

MEDLINE=21074935; PubMed=11206551; [NCBI, ExPASy, EBI, Israel, Japan]

Perna N.T., Plunkett G. III, Burland V., Mau B., Glasner J.D., Rose D.J., Mayhew G.F., Evans P.S., Gregor J., Kirkpatrick H.A., Posfai G., Hackett J., Klink S., Boutin A., Shao Y., Miller L., Grotbeck E.J., Davis N.W., Lim A., Dimalanta E.T., Potamousis K., Apodaca J., Anantharaman T.S., Lin J., Yen G., Schwartz D.C., Welch R.A., Blattner F.R.;

"Genome sequence of enterohaemorrhagic Escherichia coli O157:H7."; Nature 409:529-533(2001).

[5] SEQUENCE FROM NUCLEIC ACID.

STRAIN=0157:H7 / RIMD 0509952;

MEDLINE=21156231; PubMed=11258796; [NCBI, ExPASy, EBI, Israel, Japan]

Hayashi T., Makino K., Ohnishi M., Kurokawa K., Ishii K., Yokoyama K., Han C.-G., Ohtsubo E., Nakayama K., Murata T., Tanaka M., Tobe T., Iida T., Takami H., Honda T., Sasakawa C.,

Ogasawara N., Yasunaga T., Kuhara S., Shiba T., Hattori M., Shinagawa H.;

"Complete genome sequence of enterohemorrhagic Escherichia coli O157:H7 and genomic comparison with a laboratory strain K-12.";

DNA Res. 8:11-22(2001).

[6] SEQUENCE OF 1-71 FROM NUCLEIC ACID.

MEDLINE=93077472; PubMed=1447153; [NCBI, ExPASy, EBI, Israel, Japan]

Ueki M., Wachi M., Jung H.K., Ishino F., Matsuhashi M.;

"Escherichia coli mraR gene involved in cell growth and division.";

J. Bacteriol. 174:7841-7843(1992).

[7] SEQUENCE OF 1-41 FROM NUCLEIC ACID.

MEDLINE=93077455; PubMed=1332942; [NCBI, ExPASy, EBI, Israel, Japan]

Guzman L.M., Barondess J.J., Beckwith J.;

"FtsL, an essential cytoplasmic membrane protein involved in cell division in Escherichia coli."; J. Bacteriol. 174:7716-7728(1992).

[8] SEQUENCE OF <u>550-588</u> FROM NUCLEIC ACID.

MEDLINE=90328986; PubMed=2198024; [NCBI, ExPASy, EBI, Israel, Japan]

Michaud C., Parquet C., Flouret B., Blanot D., van Heijenoort J.;

"Revised interpretation of the sequence containing the murE gene encoding the UDP-N-acetylmuramyl-tripeptide synthetase of Escherichia coli."; Biochem. J. 269:277-280(1990).

OL MUTACENIEGIG OF GED 207

[9] MUTAGENESIS OF SER-307.

MEDLINE=86117937; PubMed=3911028; [NCBI, ExPASy, EBI, Israel, Japan]

Houba-Herin N., Hara H., Inouye M., Hirota Y.;

"Binding of penicillin to thiol-penicillin-binding protein 3 of Escherichia coli: identification of its active site.";

Mol. Gen. Genet. 201:499-504(1985).

[10] PROCESSING.

MEDLINE=90036670; PubMed=2681146; [NCBI, ExPASy, EBI, Israel, Japan]

Nagasawa H., Sakagami Y., Suzuki A., Suzuki H., Hara H., Hirota Y.;

"Determination of the cleavage site involved in C-terminal processing of penicillin-binding protein 3 of Escherichia coli.";

J. Bacteriol. 171:5890-5893(1989).

[11] MUTANT PBPBR1.

MEDLINE=89008105; PubMed=3049550; [NCBI, ExPASy, EBI, Israel, Japan] Taschner P.E., Ypenburg N., Spratt B.G., Woldringh C.L.;

"An amino acid substitution in penicillin-binding protein 3 creates pointed polar caps in Escherichia coli.":

J. Bacteriol. 170:4828-4837(1988).

[12] TOPOLOGY.

MEDLINE=90014188; PubMed=2677607; [NCBI, ExPASy, EBI, Israel, Japan] Bowler L.D., Spratt B.G.;

"Membrane topology of penicillin-binding protein 3 of Escherichia coli."; Mol. Microbiol. 3:1277-1286(1989).

Comments

- FUNCTION: CELL WALL FORMATION. ESSENTIAL FOR THE FORMATION OF A SEPTUM OF THE MUREIN SACCULUS. SYNTHESIS OF CROSS-LINKED PEPTIDOGLYCAN FROM THE LIPID INTERMEDIATES.
- PATHWAY: FINAL STAGES IN PEPTIDOGLYCAN SYNTHESIS.
- SUBCELLULAR LOCATION: INNER MEMBRANE. THE BULK OF THE MOLECULE, EXCEPT FOR THE N-TERMINAL MEMBRANE ANCHOR REGION, PROTRUDES INTO THE PERIPLASMIC SPACE, WHERE IT ACTS ON MUREIN.
- **DOMAIN**: HAS AN N-TERMINAL PENICILLIN INSENSITIVE TRANSGLYCOSYLASE DOMAIN (FORMATION OF LINEAR GLYCAN STRANDS) & A CARBOXY-TERMINAL PENICILLIN-SENSITIVE TRANSPEPTIDASE DOMAIN (CROSS-LINKING OF THE PEPTIDE SUBUNITS).
- *MISCELLANEOUS*: MUTANT PBPBR1 WHICH WAS OBTAINED AFTER SELECTION FOR INCREASED RESISTANCE TO CEPHALEXIN, CAUSES A CHANGE IN THE SHAPE OF THE CELL: THE POLAR CAPS ARE POINTED.
- SIMILARITY: BELONGS TO THE TRANSPEPTIDASE FAMILY.

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Cross-references

	K00137; AAA24300.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	X55034; CAA38861.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	D10483; BAB96652.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	AE000118; AAC73195.1;[EMBL / GenBank / DDBJ] [CoDingSequence]
EMBL	AE005185; AAG54388.1;[EMBL / GenBank / DDBJ] [CoDingSequence]
	AP002550; BAB33511.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	S49802; AAB24312.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	S49875; AAB24310.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	X55814; CAA39333.1; [EMBL / GenBank / DDBJ] [CoDingSequence]
	A03419; ZPECP3.
DID	D45070 D45070

PIR B45278; B45278.

S40594; S40594.

MEROPS S12.UNW; -. EcoGene EG10341; ftsI. EcoCyc EG10341; ftsI.

CMR P04286; B0084. IPR005311; PBP dimer. IPR001460; Transpeptdse. InterPro Graphical view of domain structure. PF00905; Transpeptidase; 1. Pfam PF03717; PBP_dimer; 1. **ProDom** [Domain structure / List of seq. sharing at least 1 domain]. **BLOCKS** P04286. P04286. ProtoNet ProtoMap P04286. **PRESAGE** P04286. DIP P04286. ModBase P04286.

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Keywords

Inner membrane; Peptidoglycan synthesis; Cell division; Cell wall; Antibiotic resistance; Multifunctional enzyme; Cell shape; Transmembrane; Complete proteome.

Features

Key	From	To	Length	Description
CHAIN	1	577	577	PEPTIDOGLYCAN SYNTHETASE
				FTSI.
PROPEP	578	588	11	
DOMAIN	1	18	18	CYTOPLASMIC (POTENTIAL).
TRANSMEM	19	39	21	POTENTIAL.
DOMAIN	40	<u> 577</u>	538	PERIPLASMIC (POTENTIAL).
ACT_SITE	307	307		ACYLATED BY PENICILLIN.
VARIANT	361	361		N -> S (IN MUTANT PBPBR1).
MUTAGEN	307	307		S->A,T: UNABLE TO BIND
MUTAGEN	307	307		PENICILLIN. S->C: STILL ABLE TO BIND PENICILLIN.



Feature aligner



Feature table viewer

Sequence information

	AA [This is th unprocessed	[This is th	weight: 6387 the MW of the ed precursor]	CR		3D5980B2CD [This the sequence]
10	20	30	40	50	60	
					1	
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70	80	90	100	110	120	
		1	1		1	
SLRVQQVSTS	RGMITDRSGR	PLAVSVPVKA	IWADPKEVHD	AGGISVGDRW	KALANALNIP	
130	140	150	160	170	180	
	1	1	1	i	1	
LDQLSARINA	NPKGRFIYLA	RQVNPDMADY	IKKLKLPGIH	LREESRRYYP	SGEVTAHLIG	
190	200	210	220	230	240	
	1	1	1	1	1	

FTNVDSQGIE GVEKSFDKWL TGQPGERIVR KDRYGRVIED ISSTDSQAAH NLALSIDERL

250	260	270	280	290	300	
I				I		
QALVYRELNN	AVAFNKAESG	SAVLVDVNTG	EVLAMANSPS	YNPNNLSGTP	KEAMRNRTIT	
310 I DVFEPGSTVK	1	330 GVVRENSVLN	340 TIPYRINGHE	350 IKDVARYSEL	360 TLTGVLQKSS	
370	380	390	400	410	420	
NVGVSKLALA	MPSSALVDTY	SRFGLGKATN	LGLVGERSGL	YPQKQRWSDI	ERATFSFGYG	
430 LMVTPLQLAR	1	450 YRPLSITKVD	460 I PPVPGERVFP	470 ESIVRTVVHM	480 MESVALPGGG	
490	500	510	520	530	540	
I						
GVKAAIKGYR	IAIKTGTAKK	VGPDGRYINK	YIAYTAGVAP	ASQPRFALVV	VINDPQAGKY	
550 I YGGAVSAPVF	560 GAIMGGVLRT	570 MNIEPDALTT	580 GDKNEFVINQ	GEGTGGRS		P04286 in <u>FASTA</u> format

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Direct BLAST submission at NCBI (Bethesda, USA)



ScanProsite, MotifScan



Sequence analysis tools: <u>ProtParam</u>, <u>ProtScale</u>, <u>Compute pI/Mw</u>, <u>PeptideMass</u>, <u>PeptideCutter</u>, <u>Dotlet</u> (Java)



Feature table viewer (Java)



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